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(54) Title: METHOD FOR REDUCING C-RECTIVE PROTEIN LEVELS WITH NON-ANTIBACTERIAL TETRACYCLINE **FORMULATIONS**

(57) Abstract: The present invention is for a method for decreasing C-reactive protein levels (CRP) in a mammal in need thereof. The method comprises administering an effective amount of a non-antibacterial tetracycline formulation, to the mammal. In one embodiment, the non-antibacterial tetracycline formulation is a non-antibacterial amount of an antibacterial tetracycline. In another embodiment, the non-antibacterial tetracycline formulation is a non-antibacterial tetracycline.

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METHOD FOR REDUCING C-REACTIVE PROTEIN LEVELS WITH NON-ANTIBACTERIAL TETRACYCLINE FORMULATIONS

BACKGROUND OF THE INVENTION

C-reactive protein (CRP) is a special type of protein referred to as an acute phase reactant. Acute phase reactants, such as CRP, are released by the body in response to acute injury, infection or other inflammatory conditions, such as, for example, atherosclerosis.

Atherosclerosis is a condition in which atheromatous plaques form in the arteries. Atheromatous plaques are deposits, or degenerative accumulations, of lipids on the innermost layer of the wall of an artery. Such plaques contain inflammatory cells. The rupture of atheromatous plaques is thought to be the mechanism for acute myocardial infarction (e.g. heart attack).

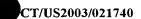
The release of acute phase reactants, such as CRP, in response to inflammation, has been proposed as a potential marker of coronary artery diseases, due to, for example, atherosclerosis. Accordingly, current research is focusing on developing drugs that inhibit CRP, and thus, decrease the incidence of such diseases. See, Taubes, Gary, *Does Inflammation Cut to the Heart of the Matter?*, Science, 12 April 2002; 296: 242-245. For example, recent studies have shown that treatment with pravastatin, an HMG-CoA reductase inhibitor (i.e. statin), appears to result in reduced levels of CRP. Ridker, P., Nader, R., et al. *Long Term Effects of Pravastin on Plasma Concentration of C-Reactive Protein*, Circulation, 1999;100:230-235.

However, HMG-CoA reductase inhibitors, such as pravastatin, are associated with numerous side effects. These side effects include constipation, stomach pain, nausea and vomiting.

Therefore, the prior art treatments for reducing CRP levels are limited and not without adverse effects. There is a need for novel, alternate, and superior treatments for reducing CRP levels.

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The compound tetracycline is a member of a class of antibiotic compounds that is referred to as the tetracyclines, tetracycline compounds, tetracycline derivatives and the like. The compound tetracycline exhibits the following general structure:

5 The numbering system of the tetracycline ring nucleus is as follows:

Structure B

Tetracycline, as well as the terramycin and aureomycin derivatives, exist in nature, and are well known antibiotics. Natural tetracyclines may be modified without losing their antibiotic properties, although certain elements must be retained. The modifications that may and may not be made to the basic tetracycline structure have been reviewed by Mitscher in *The Chemistry of Tetracyclines*, Chapter 6, Marcel Dekker, Publishers, New York (1978). According to Mitscher, the substituents at positions 5-9 of the tetracycline ring system may be modified without the complete loss of antibiotic properties.

Changes to the basic ring system or replacement of the substituents at positions 4 and 10-12a, however, generally lead to synthetic tetracyclines with substantially less or effectively no antimicrobial activity. Some examples of chemically modified non-antibacterial tetracyclines (hereinafter CMTs) are 4-dedimethylaminotetracyline, 4-dedimethylaminosancycline (6-demethyl-6-deoxy-4-dedimethylaminotetracycline), 4-dedimethylaminominocycline (7-dimethylamino-6-

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demethyl-6-deoxy-4-dedimethylaminotetracycline), and 4-dedimethylaminodoxycycline (5-hydroxy-6-deoxy-4-dedimethylaminotetracycline).

In addition to their antimicrobial properties, tetracyclines have been described as having a number of other uses. For example, tetracyclines are also known to inhibit the activity of collagen destructive enzymes produced by mammalian 5 (including human) cells and tissues by non-antibiotic mechanisms. Such enzymes include the matrix metalloproteinases (MMPs), including collagenases (MMP-1, MMP-8 and MMP-13), gelatinases (MMP-2 and MMP-9), and others (e.g. MMP-12, MMP-14). See Golub et al., J. Periodont. Res. 20:12-23 (1985); Golub et al. Crit. Revs. Oral Biol. Med. 2:297-322 (1991); U.S. Patent Nos. 4,666,897; 4,704,383; 10 4,935,411; 4,9354,412. Also, tetracyclines have been known to inhibit wasting and protein degradation in mammalian skeletal muscle, U.S. Pat. No. 5,045,538, to inhibit inducible NO synthase, U.S. Patent Nos. 6,043,231 and 5,523,297, and phospholipase A2, U.S. Patent Nos. 5,789,395 and 5,919,775, and to enhance IL-10 production in mammalian cells. These properties cause the tetracyclines to be useful in treating a 15 number of diseases.

The object of this invention is to provide a method for reducing C-reactive protein levels in a mammal in need thereof.

SUMMARY OF THE INVENTION

It has now been discovered that this and other objectives can be achieved by the present invention. The present invention provides a method for decreasing C-reactive protein levels (CRP) in a mammal in need thereof. The method comprises administering an effective amount of a non-antibacterial tetracycline formulation, to the mammal.

In one embodiment, the non-antibacterial tetracycline formulation is a nonantibacterial amount of an antibacterial tetracycline. In another embodiment, the nonantibacterial tetracycline formulation is a non-antibacterial tetracycline.

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DETAILED DESCRIPTION

The invention relates to decreasing C-reactive protein levels by administering a non-antibacterial tetracycline formulation.

In one embodiment of the invention, the non-antibacterial tetracycline formulation is an antibacterial tetracycline compound administered in a non-antibacterial amount, as will be discussed below. For this embodiment, the tetracycline may be any such tetracycline having clinically significant antibacterial activity.

Some examples of antibacterial tetracyclines include tetracycline, as well as the 5-OH (oxytetracycline, e.g. Terramycin) and 7-Cl (chlorotetracycline, e.g. Aureomycin) derivatives, which exist in nature. Semi-synthetic tetracyclines, which include, for example, doxycycline, minocycline and sancycline, can also be used for this embodiment. Examples also include demeclocycline and lymecycline.

In another embodiment of the invention, the non-antibacterial tetracycline formulation is a non-antibacterial tetracycline compound. Non-antibiotic tetracycline compounds are structurally related to the antibiotic tetracyclines, but have had their antibiotic activity substantially or completely eliminated by chemical modification, as mentioned above. For example, non-antibiotic tetracycline compounds are incapable of achieving antibiotic activity comparable to that of doxycycline unless the concentration of the non-antibiotic tetracycline is at least about ten times, preferably at least about twenty five times, greater than that of doxycycline.

One such group of chemically modified non-antibacterial tetracyclines (CMT's) includes any of the 4-dedimethylaminotetracycline derivatives, for example, 4-dedimethylaminosancycline (CMT-3), 4-dedimethylaminodoxycycline (CMT-8) and 4-dedimethylaminominocycline (CMT-10).

Some additional examples of generic and specific chemically modified, non-antibiotic tetracycline compounds that are suitable for use in the method of the invention are found in PCT/US01/16272. All such generic and specific compounds are incorporated herein by reference.

Some preferred examples of suitable 4-dedimethylaminotetracycline derivatives include the following general formulae (I) through (IV):

General Formula (I)

Structure A represents the 4-dedimethylaminosancycline (CMT-3) derivatives

Structure A

wherein R7, R8, and R9 taken together in each case, have the following meanings:

	R7	R8	<u>R9</u>
	azido	hydrogen	hydrogen
10	dimethylamino	hydrogen	azido
	hydrogen	hydrogen	azido
	dimethylamino	hydrogen	amino
	acylamino	hydrogen	hydrogen
	amino	hydrogen	nitro
15	hydrogen	hydrogen (N,Ndimethyl)	glycylamino
	amino	hydrogen	amino
	hydrogen	hydrogen ethoxythio	carbonylthio
	dimethylamino	hydrogen	acylamino
	dimethylamino	hydrogen	diazonium
20	dimethylamino	chloro	amino
	hydrogen	chloro	amino
	amino	chloro	amino
	acylamino	chloro	acylamino

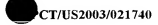
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		amino		chloro	hydrogen
		acylamino		chloro	hydrogen
		monoalkylan	nino	chloro	amino
		nitro		chloro	amino
5		dimethylami	no	chloro	acylamino
		dimethylami	no	chloro	dimethylamino
		acylamino		hydrogen	hydrogen
		hydrogen		hydrogen	acylamino
	(CMT-301)	bromo		hydrogen	hydrogen
10	(CMT-302)	nitro		hydrogen	hydrogen
	(CMT-303)	hydrogen		hydrogen	nitro
	(CMT-304)	acetamido		hydrogen	hydrogen
	(CMT-305)	hydrogen		hydrogen	acetamido
	(CMT-306)	hydrogen		hydrogen	dimethylamino
15	(CMT-307)	amino		hydrogen	hydrogen
	(CMT-308)	hydrogen		hydrogen	amino
	(CMT-309)	hydrogen		hydrogen	dimethylaminoacetamido
	(CMT-310)	dimethylamir	10	hydrogen	hydrogen
	(CMT-311)	hydrogen		hydrogen	palmitamide
20					
		<u>R7</u>	R8	R9	R2
	(CMT-312)	hydrogen	hydrogen	hydrogen	CONHCH ₂ -pyrrolidin-1-yl
	(CMT-313)	hydrogen	hydrogen	hydrogen	CONHCH ₂ -piperadin-1-yl
25	(CMT-314)	hydrogen	hydrogen	hydrogen	CONHCH ₂ -morpholin-1-yl
,	(CMT-315)	hydrogen	hydrogen	hydrogen	CONHCH2-piperazin-1-yl

General Formula (II)

Structures B through E represent the 4-dedimethylaminodoxycycline (CMT-8) derivatives

5 wherein R7, R8, and R9 taken together in each case, have the following meanings:

	<u>R7</u>	R8	<u>R9</u>
	azido	hydrogen	hydrogen
	dimethylamino	hydrogen	azido
	hydrogen	hydrogen	azido
10	dimethylamino	hydrogen	amino



				1 1	1 1
		acylamino		hydrogen	hydrogen
	•	hydrogen		hydrogen	acylamino
		amino		hydrogen	nitro
		hydrogen		hydrogen	(N,N-dimethyl)glycylamino
5		amino		hydrogen	amino
		hydrogen		hydrogen	ethoxythiocarbonylthio
		dimethylamir	10	hydrogen	acylamino
		hydrogen		hydrogen	diazonium
		diazonium		hydrogen	hydrogen
10		ethoxythioca	bonylthio	hydrogen	hydrogen
		dimethylamir	10	chloro	amino
		amino		chloro	amino
		acylamino		chloro	acylamino
		hydrogen		chloro	amino
15		amino		chloro	hydrogen
		acylamino		chloro	hydrogen
		monoalkylam	nino	chloro	amino
		nitro		chloro	amino
	(CMT-801)	hydrogen	•	hydrogen	acetamido
20	(CMT-802)	hydrogen		hydrogen	dimethylaminoacetamido
	(CMT-803)	hydrogen	1	hydrogen	palmitamide
	(CMT-804)	hydrogen		hydrogen	nitro
	(CMT-805)	hydrogen		hydrogen	amino
	(CMT-806)	hydrogen		hydrogen	dimethylamino
25	,	, .			·
		R7	R8	R9	R2
	(CMT-807)	hydrogen	hydrogen	hydrogen	CONHCH ₂ -pyrrolidin-1-yl
	(CMT-808)	hydrogen	hydrogen	hydrogen	CONHCH ₂ -piperadin-1-yl
	(CMT-809)	hydrogen	hydrogen	hydrogen	CONHCH ₂ -piperazine-1-yl
	(CIVII -007)	, 0 50	, 0 50.11	, 55	

General Formula (III)

Structure F represents the 4-dedimethylaminominocycline (CMT-10) derivatives

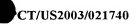
Structure F

wherein R8 is hydrogen or halogen and R9 is selected from the group consisting of nitro (CMT-1002), (N,N-dimethyl)glycylamino, ethoxythiocarbonylthio. A compound related to structure F has a 7-trimethylammonium group instead of the 7-diemthylamino group, i.e. 7-trimethylammoniumsancycline (CMT-1001), and

General Formula (IV)

Structure G

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wherein R7, R8, and R9 taken together in each case, have the following meanings:

	7.5		C.2
	<u>R7</u>	R8	<u>R9</u>
	amino	hydrogen	hydrogen
5	nitro	hydrogen	hydrogen
	azido	hydrogen	hydrogen
	dimethylamino	hydrogen	azido
	hydrogen	hydrogen	amino
	hydrogen	hydrogen	azido
10	hydrogen	hydrogen	nitro
	bromo	hydrogen	hydrogen
	dimethylamino	hydrogen	amino
	acylamino	hydrogen	hydrogen
	hydrogen	hydrogen	acylamino
15	amino	hydrogen	nitro
	hydrogen	hydrogen	(N,N-dimethyl)glycylamino
1	amino	hydrogen	amino
·	diethylamino	hydrogen	hydrogen
	hydrogen	hydrogen	ethoxythiocarbonylthio
20	dimethylamino	hydrogen	methylamino
	dimethylamino	hydrogen	acylamino
	dimethylamino	chloro	amino
	amino	chloro	amino
	acylamino	chloro	acylamino
25	hydrogen	chloro	amino
	amino	chloro	hydrogen
	acylamino	chloro	hydrogen

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monoalkylamino nitro chloro chloro amino amino

Additional CMT's for purposes of the invention include, 4-dedimethylaminotetracycline (CMT-1), tetracycline nitrile (CMT-2), 4-dedimethylaminochlorotetracycline (CMT-4), 4-dedimethylamino-4-hydroxytetracycline (CMT-6), 2a-dehydroxy-4-dedimethylaminotetracycline (CMT-7), and 1-deoxy-12a-dehydroxy-4-dedimethylaminotetracycline (CMT-9).

The chemically modified tetracyclines can be made by methods known in the art. See, for example, Mitscher, L.A., *The Chemistry of the Tetracycline Antibiotics*, Marcel Dekker, New York (1978), Ch. 6, and U.S. Patents 4,704,383 and 5,532,227.

The invention also includes pharmaceutically acceptable salts of the above disclosed compounds. The present invention embraces salts, including acid-addition and metal salts of the 4-dedimethylaminotetracycline compounds described herein. Such salts are formed by well known procedures. By "pharmaceutically acceptable salts" is meant salts that do not substantially contribute to the toxicity of the compound.

Some examples of suitable salts include salts of basic tetracycline compounds and mineral acids such as hydrochloric, hydriodic, hydrobromic, phosphoric, metaphosphoric, nitric and sulfuric acids, as well as salts of organic acids such as tartaric, acetic, citric, malic, benzoic, glycollic, gluconic, gulonic, succinic, arylsulfonic, e.g. p-toluenesulfonic acids, and the like.

After preparation, the novel compounds of the present invention can be conveniently purified by standard methods known in the art. Some suitable examples include crystallization from a suitable solvent or partition-column chromatography.

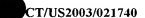
The preferred pharmaceutical composition for use in the method of the invention includes a combination of the tetracycline compound in a suitable pharmaceutical carrier (vehicle) or excipient as understood by practitioners in the art. Examples of carriers and excipients include starch, milk, sugar, certain types of clay,

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gelatin, stearic acid or salts thereof, magnesium or calcium stearate, talc, vegetable fats or oils, gums and glycols.

The tetracycline compounds of the invention may be administered by methods known in the art, typically, systemically. Systemic administration can be enteral or parenteral. Enteral administration is a preferred route of delivery of the tetracycline, and compositions including the tetracycline compound with appropriate diluents, carriers, and the like are readily formulated. Liquid or solid (e.g., tablets, gelatin capsules) formulations can be employed.

Administration can also be accomplished by a nebulizer or liquid mist.

Nebulization is a preferred route of delivery of the tetracycline in situations where the respiratory system is particularly infected. By utilizing a nebulizer, the tetracycline is taken directly into the individuals respiratory system through inspiration.

Parenteral administration of the tetracycline compounds of the invention (e.g., intravenous, intramuscular, subcutaneous injection) is also contemplated.

Formulations using conventional diluents, carriers, etc. such as are known in the art can be employed to deliver the compound.

The tetracycline compound may be administered to mammals by sustained release, as is known in the art. Sustained release administration is a method of drug delivery to achieve a certain level of the drug over a particular period of time.

The amount of tetracycline compound administered is any amount effective for decreasing CRP levels in the mammal in need thereof. The actual preferred amounts of tetracycline compound in a specified case will vary according to the particular compositions formulated, the mode of application, and the particular subject being treated. The appropriate dose of the tetracycline compound can readily be determined by those skilled in the art.

The minimal amount of the tetracycline compound administered to a human is the lowest amount capable of providing effective treatment of the conditions.

Effective treatment is a decrease in CRP levels, of the mammal.

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The maximal amount for a mammal is the highest amount that does not cause undesirable or intolerable side effects. Such doses can be readily determined by those skilled in the art.

The amount of an antibacterial tetracycline is an amount that has substantially no antibacterial activity, i.e. an amount that does not significantly prevent the growth of bacteria. For example, tetracycline compounds that have significant antibacterial activity may be administered in an amount which is 10-80% of the antibacterial amount. More preferably, the antibacterial tetracycline compound is administered in an amount which is 40-70% of the antibacterial amount.

The amount of tetracycline administered may be measured, for example, by a daily dose or by serum level. Some examples of non-antibiotic daily doses of antibiotic tetracyclines, based on steady-state pharmacokinetics, are as follows: 20 mg/twice a day for doxycycline; 38 mg of minocycline one, two, three or four times a day; 60 mg of tetracycline one, two, three or four times a day, 1000mg/day of oxytetracycline, 600mg/day of demeclocycline and 600mg/day of lymecycline.

In a preferred embodiment, doxycycline is administered in a daily amount of from about 10 to about 60 milligrams, preferably 30 to 60 milligrams, but maintains a concentration in human plasma below the threshold for a significant antibiotic effect.

In an especially preferred embodiment, doxycycline hyclate is administered at a 20 milligram dose twice daily. Such a formulation is sold for the treatment of periodontal disease by CollaGenex Pharmaceuticals, Inc. of Newtown, Pennsylvania under the trademark Periostat ®.

Antibiotic serum levels are also known in the art. For example, a single dose of two 100 mg minocycline HCl tablets administered to an adult human results in minocycline serum levels ranging from 0.74 to 4.45 μ g/ml over a period of an hour. The average level is 2.24 μ g/ml.

Two hundred and fifty milligrams of tetracycline HCl administered every six hours over a twenty-four hour period produces a peak plasma concentration of approximately 3 µg/ml. Five hundred milligrams of tetracycline HCl administered

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every six hours over a twenty-four hour period produces a serum concentration level of 4 to 5 µg/ml.

In general, the tetracycline compound is administered in an amount which results in a serum concentration between about 0.1 and 10.0 μ g/ml, more preferably between 0.3 and 5.0 μ g/ml. For example, doxycycline, in a non-antibacterial formulation, is administered in an amount which results in a serum concentration between about 0.1 and 0.8 μ g/ml, more preferably between 0.4 and 0.7 μ g/ml.

Non-antibacterial tetracycline compounds can be used in higher amounts than antibacterial tetracyclines, while avoiding the indiscriminate killing of bacteria, and the emergence of resistant bacteria. For example, 6-demethyl-6-deoxy-4-dedimethylaminotetracycline (CMT-3) may be administered in doses of about 10 to about 200mg/day, or in amounts that result in serum levels in humans of about 1.0µg/ml to about 10µg/ml. For example, a dose of about 10 to about 20mg/day produces serum levels in humans of about 1.0 µg/ml.

For example, CMTs can be systemically administered in a mammal in a minimal amount of about 0.05mg/kg/day to about 0.3mg/kg/day, and a maximal amount of about 18mg/kg/day to about 60mg/kg/day. The practitioner is guided by skill and knowledge in the field, and the present invention includes, without limitation, dosages that are effective to achieve the desired antibacterial activity.

The tetracyclines of the present invention decrease CRP levels in mammals in need thereof. CRP, as discussed above, is a special type of protein produced during inflammation.

A mammal in need of decreasing CRP levels is any mammal that has an elevated CRP level. For example, a mammal having a condition associated with inflammation will have an elevated CRP level. Conditions associated with inflammation include, for example, cardiac conditions, cerebrovascular disease, arthritis, asthma, periodontitis, cancer, and lupus.

Cardiovascular conditions include, for example, myocardial infarction, atherosclerosis, and angina. Cerebrovascular disease includes stroke and aneurysm.

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A mammal which can benefit from the methods of the present invention could be any mammal. Categories of mammals include, for example, humans, farm animals, domestic animals, laboratory animals, etc. Some examples of farm animals include cows, pigs, horses, goats, etc. Some examples of domestic animals include dogs, cats, etc. Some examples of laboratory animals include rats, mice, rabbits, guinea pigs, etc.

Examples

The following exemplary data serves to provide further appreciation of the invention but are not meant in any way to restrict the effective scope of the invention.

A prospective, randomized study was conducted over six (6) months to investigate the effectiveness of low-dose doxycycline (LDD) versus placebo, in the prevention of subsequent plaque rupture events in patients enrolled after an initial acute coronary syndrome.

Biochemical markers of inflammation were assessed at study entry and after six (6) months of therapy in a subset of patients. A total of thirty (30) patients completed the study of whom thirteen (13) were randomized to placebo and seventeen (17) to LDD. There were no significant differences in age, male gender, hypertension, diabetes, smoking, previous cardiac history, extent of coronary disease, presentation with acute myocardial infarction or unstable angina, or percutaneous coronary intervention between LDD and placebo treated patients.

At six months clinical follow-up, there was no difference in the composite endpoint of cardiovascular death, myocardial infarction or troponin-positive unstable angina in LDD compared to placebo treated patients. As demonstrated in Table I, C-reactive protein (CRP) levels were reduced by 46% from 4.8µg/ml to 2.6µg/ml (P<0.05) among patients randomized to LDD. In placebo-treated patients, CRP was 5.2 µg/ml at study entry and 4.9 µg/ml at six months (P=not significant (n.s.)).

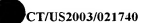


Table I

The Effect of Low-Dose Doxycycline (LDD) on C-Reactive Protein in

Patients With Acute Myocardial Syndrome: Preliminary Data¹

5		Placebo (n=13 subjects)	Low-Dose Doxycycline (n=17 subjects)
	Baseline CRP level	$5.2 \pm 0.8 \ \mu g/ml$	$4.8 \pm 0.6 \mu \text{g/ml}$
10	6-Month CRP level	$4.9 \pm 0.7^2 \mu \text{g/ml}$	$2.6 \pm 0.4^3 \mu \text{g/ml}$
15	Reduction Due To Treatment	5%	45%

¹ Each value represents the mean ± S.E.M.

Table II demonstrates the preferential efficacy of LDD at decreasing CRP levels in patients having higher baseline CRP levels. LDD-treated patients having lower baseline CRP levels showed a 23% reduction (3.0 μg/ml to 2.3 μg/ml (P= n.s.)). Patients with higher baseline CRP levels were reduced by 58% from 7.2 μg/ml to 3.0 μg/ml (P<0.001). In placebo-treated patients with higher baseline CRP, CRP levels were decreased by 23% (7.1 μg/ml to 5.5 μg/ml).

² not significant (n.s.) comparing six-month values to baseline values.

³ P<0.05



Table II

Low Dose Doxycycline (LDD) Preferentially Suppresses C-Reactive Protein in

Patients With Higher CRP Values at Baseline

5	Patients with lower baseline CRP (≤5 μg/ml)			Patients with higher CRP (>5 µg/ml)	
		Placebo	LDD	Placebo	LDD
10	Baseline CRP	2.9 ± 0.6	3.0 ± 0.4	7.1 ± 0.8	7.2 ± 0.6
	Six Month CRP	4.2 ± 0.9^{b}	2.3 ± 0.5^{b}	5.5 ± 1.1^{b}	$3.0\pm0.7^{\rm a}$
15	Reduction Due To Treatment		23%	23%	58%
13	^a P < 0.001				

b n.s.

We Claim:

- 1. A method for decreasing C-reactive protein levels in a mammal in need thereof, the method comprising administering to the mammal an effective amount of a non-antibacterial tetracycline formulation.
- 2. The method according to claim 1, wherein the tetracycline formulation comprises a non-antibacterial amount of an antibacterial tetracycline.
- 3. The method according to claim 2, wherein the antibacterial tetracycline is selected from the group consisting of terramycin, aureomycin, doxycycline, minocycline, tetracycline, oxytetracycline, chlortetracycline, demeclocycline, lymecycline, or pharmaceutically acceptable salts thereof.
- 4. The method according to claim 1, wherein the tetracycline formulation comprises a non-antibacterial tetracycline
- 5. The method of claim 4, wherein the non-antibacterial tetracycline is selected from the group consisting of CMT-1, CMT-2, CMT-4, CMT-6, CMT-7 or CMT-9, or pharmaceutically acceptable salts thereof.
- 6. The method of claim 4, wherein the tetracycline is CMT-3, or its analogs, or pharmaceutically acceptable salts thereof.
- 7. The method according to claim 4, wherein the tetracycline is CMT-8, or its analogs, or pharmaceutically acceptable salts thereof.
- 8. The method according to claim 4, wherein the tetracycline is CMT-10, or its analogs, or pharmaceutically acceptable salts thereof.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US03/21740

	SIFICATION OF SUBJECT MATTER					
IPC(7) US CL	: A61K 31/65 : 514/152					
	International Patent Classification (IPC) or to both na	tional classi	fication and IPC	·····		
	DS SEARCHED					
Minimum doc U.S.: 51	umentation searched (classification system followed b 4/152	y classifica	tion symbols)			
Documentation NONE	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched NONE					
Electronic data WEST	a base consulted during the international search (name	e of data ba	se and, where practicable, searc	ch terms used)		
C. DOCU	IMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where a			Relevant to claim No.		
A	US 4,666,897 A (GOLUB et al.) 19 May 1987(19.0)	5.1987), sec	e entire document.	1-8		
	documents are listed in the continuation of Box C.		See patent family annex.	national filing days as associate		
"A" document	defining the general state of the art which is not considered to be ar relevance	1	date and not in conflict with the applica principle or theory underlying the inven	tion but cited to understand the		
"E" earlier app	olication or patent published on or after the international filing date	"X"	document of particular relevance; the cl considered novel or cannot be considered when the document is taken alone			
	which may throw doubts on priority claim(s) or which is cited to he publication date of another citation or other special reason (as	"Y"	document of particular relevance; the cl considered to involve an inventive step combined with one or more other such	when the document is		
"O" document	referring to an oral disclosure, use, exhibition or other means		being obvious to a person skilled in the			
	"P?" document published prior to the international filing date but later than the "&" document member of the same patent family priority date claimed					
Date of the ac	Date of the actual completion of the international search Date of mailing of the international search Date of mailing of the international search					
28 November 2003 (28.11.2003)						
Name and mailing address of the ISA/US Mail Stop PCT, Attn: ISA/US Commissioner for Patents P.O. Box 1450 Alexandria, Virginia 22313-1450 Facsimile No. (703)305-3230 Authorized officer Marianne Seidel Telephone No. (703) 308-0196						

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